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The Blood Exposome and Its Role in Discovering Causes of Disease

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Abstract

Background. Since 2001 researchers have mainly examined the human genome (G) to discover causes of disease despite evidence that G explains relatively little risk. We posit that unexplained disease risks are caused by the exposome (E, representing all exposures) and G×E interactions. It follows that etiologic research has been hampered by scientists' continuing reliance on low-tech methods to characterize E as contrasted with high-tech omics for characterizing G.

Objectives: Because exposures are inherently chemical in nature and arise from both endogenous and exogenous sources, blood specimens can be used to characterize exposomes. To explore the 'blood exposome' and its connection to disease we sought human-blood concentrations of many chemicals along with their sources, evidence of chronic-disease risks and numbers of metabolic pathways.

Methods: From the literature we obtained human-blood concentrations for 1,561 small molecules and metals, derived from foods, drugs, pollutants and endogenous processes. Chemical similarities were mapped after weighting by blood concentrations, disease-risk citations and numbers of human metabolic pathways.

Results: Blood concentrations spanned 11 orders of magnitude and were indistinguishable for endogenous and food chemicals and drugs while those of pollutants were 1,000-times lower. Chemical similarities mapped by disease risks were equally distributed by source categories while those mapped by metabolic pathways were dominated by endogenous molecules and essential nutrients.

Conclusions: The complexity of human exposures motivates characterization of the blood exposome, which includes all biologically active chemicals, for studies of disease etiology. Because most small molecules in blood are not human metabolites, investigations of causal pathways should expand beyond the endogenous metabolome.

Introduction

World-wide mortality is dominated by non-communicable diseases, particularly cardiovascular disease (29%), cancer (15%) and respiratory diseases (7%) (Lozano et al. 2012). These chronic diseases result from the combined effects of the human genome (G) and exposome (E, representing all exposures). (Although geneticists use the term 'environment' to denote nongenetic factors, many scientists and the general public equate 'environment' with 'pollution', which represents only one class of exposures. We use the term 'exposome' to encompass all exogenous and endogenous exposures.) But attribution of risks to G. E and their interaction (G×E) has been problematic because of disparities in characterizing genes and exposures (Rappaport and Smith 2010; Wild 2005). In fact, sequencing the human genome in 2001 permitted researchers to comprehensively explore G and its progeny (i.e. genome \rightarrow transcriptome \rightarrow proteome \rightarrow metabolome) but did not promote detailed characterization of E, which in epidemiological and clinical research still relies on questionnaires, geographical information and targeted surveys (Ezzati and Riboli 2013; Lim et al. 2012). Also, the study of external and internal exposures (including endogenous chemicals) has focused on a limited number of molecules and metals that cannot compare with the resolution of genome-wideassociation studies (GWAS).

Interestingly, the variation in chronic-disease incidence explained by scores of GWAS has been so small that searches are underway for 'missing heritability' (Goldstein 2009; Manolio et al. 2009) and 'genetic dark matter' (Galvan et al. 2010; Martin and Chang 2012; Melhem and Devlin 2010). Even assuming that a host of rare alleles account for some unexplained phenotypic variation (Kraft and Hunter 2009), it is reasonable to posit that E and G×E are the primary causes of chronic diseases, as suggested by studies of families and twins (Hemminki et al. 2006; Lichtenstein et al. 2000), epigenetics (Gluckman et al. 2008; Gluckman et al. 2010; Smith and

Meissner 2013) and gene-expression profiles that change with lifestyles and infections (Chen et al. 2012; Preininger et al. 2013). In fact, as shown in Figure 1, half of the 50 million global deaths in 2010 were attributed to a small set of exposures, dominated by particulate air pollution (combined effects of ambient particles and household smoke), smoking (active and passive) and diet (Lim et al. 2012). This conundrum - where scientists use high-tech omics to detect small effects of G but rely upon low-tech methods to study potentially large effects of E and G×E - has produced a very uneven record of etiologic research.

One way to level the playing field would be to explore health impacts of E and G×E with exposome-wide association studies (EWAS) (Rappaport 2012) that obtain comprehensive, quantitative measurements of chemicals in human biospecimens (Holmes et al. 2008; Ritchie et al. 2010; Z Wang et al. 2011). This approach recognizes that meaningful exposures are mediated in the internal chemical environment (Rappaport and Smith 2010) by endogenous signaling molecules, exogenous chemicals and reactive electrophiles (E-factors) that communicate with cells, tissues and organs via mutations, post-translational modifications, enzymes, transcription factors and receptors (G-factors) (Brodsky and Medzhitov 2009; Liebler 2008; Menon and Manning 2013). Because blood transports chemicals to and from tissues and represents a reservoir of all endogenous and exogenous chemicals in the body at a given time (Nicholson et al. 2012b), the blood exposome offers a parsimonious but essentially unexplored means for interrogating biologically-relevant exposures (Rappaport 2012).

Methods

Sources of data

To investigate the portion of the blood exposome represented by small molecules and metals, we obtained blood concentrations of 1,561 chemicals from samples of healthy human populations compiled by the Human Metabolome Database (HMDB 2013) (Wishart et al. 2013) (1,451

chemicals) and the U.S. National Health and Nutrition Examination Survey (NHANES) (CDC 2009, 2012, 2013) (110 chemicals). Each molecule or metal was assigned one of the following four source categories: endogenous chemical (from intrinsic human metabolism, n = 1,223), food chemical (n = 195), pollutant (n = 94) or drug (n = 49). (The process for selecting chemicals is described in Supplemental Material Table S1). To link individual chemicals with chronic-disease risks and systems biology, we retrieved additional data from the U.S. National Center for Biotechnology Information (NCBI 2013) matching to citations from the PubMed database of chronic-disease-risk factors or the Biosystems database of human metabolic pathways (Biosystems 2013). Although modest in size, these samples allowed us to explore the range of human blood concentrations, to test for differences in median levels across source categories and to map chemical similarities after weighting by blood concentration, disease-risk citations and human metabolic pathways. Relevant data are given in Supplemental Material, Table S1.

HMDB entries were from metabolic studies in mostly-Western populations, and included endogenous and food chemicals, drugs and pollutants, while NHANES included only nutrients and pollutants in U.S. populations. When a given chemical was present in both databases, NHANES entries were used. If the same chemical had been reported in more than one study or year, the geometric mean concentration was used. Numbers of individual subjects varied across chemicals. Drug concentrations were reported in clinical trials at therapeutic doses.

The chemical abstract service (CAS) registry number(s) was used as the query parameter to search the PubMed database with medical subject headings (MeSH) annotations to retrieve the citations describing epidemiological studies. The search string was "(blood OR plasma OR serum) AND ("risk factors"[MeSH Terms] OR "relative risk*" OR "odds ratio*" OR "hazard ratio*")""+CAS number+"[EC/RN Number]("journal article"[pt] NOT review[pt] NOT "meta analysis"[pt]) (hasabstract[text] AND "humans"[MeSH Terms]) english[lang] (neoplasms[mesh]

OR diabetes[mesh] OR "cardiovascular diseases"[mesh] OR "Respiratory Tract Diseases"[mesh])". For retrieval of pathway hits, PubChem identifiers for each compound were searched against the NCBI Biosystems database. Chemical similarity maps were generated by Metamapp (Metamapp 2013).

Statistical analysis

Differences in median blood concentrations across source categories were evaluated with Kruskal-Wallis tests via SAS for Windows (v.9.3) (SAS Institute, Cary, NC).

Results

Blood concentrations

Cumulative distributions of blood concentrations are shown in Figure 2 for the four sources of chemicals. Concentrations ranged from 160 fM to 140 mM, a staggering 11 orders of magnitude. Within each category, concentrations covered a 10^7 -fold range. Median blood levels of endogenous chemicals (0.94 μ M), food chemicals (1.00 μ M) and drugs (0.30 μ M) were not significantly different (*P*-value = 0.246). In contrast, pollutant concentrations were 1,000 times lower (median = $2.4 \times 10^{-4} \,\mu$ M, *P*-value <0.0001) and only pollutants with blood levels above the median value overlapped with other distributions.

Chemical-similarity maps

Endogenous and dietary molecules comprised more than 100 chemical classes, particularly lipids, steroids, amino acids, fatty acids and nucleotides (Supplemental Material, Table S1). In addition to nutrients and vitamins, food chemicals included such bioactive molecules as aflatoxin-B1 (a carcinogen from mold-infected grains and nuts), solanidine (a toxin from potatoes), sulforaphane (a DNA-protective agent from cruciferous vegetables), acetaldehyde (a mutagen from metabolism of alcohol), genistein (an endocrine-disrupting chemical from soy

products) and trimethylamine-*N*-oxide (a suspected cause of atherosclerosis from metabolism of choline and carnitine). Exogenous pollutants were primarily halogenated compounds - trihalomethanes, chlorinated pesticides, perfluorinated compounds, polychlorinated biphenyls (PCBs), brominated diphenyl ethers and some chlorinated dioxins and furans - and metals, but also included a few volatile aromatic species (notably benzene) and metabolites of nicotine. This diversity is illustrated in Figure 3A, which maps the 1,561chemicals by their structural similarities (Barupal et al. 2012), with symbol sizes indicating blood concentrations. Constellations of biochemical classes were populated largely by endogenous and food chemicals, while drugs clustered with aromatic compounds (between map locations designated 'AN' and 'BD' in Figure 3A) and pollutants were mainly at map peripheries (locations 'AH' and 'AX', respectively). Metals and metalloids originated from foods (6 most abundant: Na, K, Fe, Ca, P and Mg), pollution (6 most abundant: Si, Sr, Ni, Pb, Be and As) and one drug (Li).

Since citations to risk factors summarize epidemiologic and clinical evidence associating a chemical with disease phenotypes, we found PubMed citations for 960 searchable substances in our inventory (only chemicals with CAS registry numbers were searchable in PubMed) and obtained 19,656 citations matching 336 (35%) of these chemicals. (Numbers of matching citations are included in Supplemental Material, Table S1). The distribution of citations per chemical was highly skewed, with a median value of 7.5 and a maximum of 4,499 (cholesterol). The large numbers of citations per chemical and positive skewness probably reflect publication bias in hypothesis-driven epidemiologic studies and clinical trials. Median numbers of citations varied two-fold across source categories (drugs = 10, endogenous = 6, food chemicals = 13, pollutants = 6; *P*-value = 0.041). When food chemicals were removed, median values for the other categories were not significantly different (*P*-value = 0.307). This indicates that a typical

food chemical was about twice as likely to be cited as a chronic-disease-risk factor than a chemical from another category.

The chemical-similarity map for these 336 chemicals is shown in Figure 3B, where symbol size reflects the number of citations. This map shared prominent clustering patterns with Figure 3A, except that individual lipid molecules were largely absent (lipids tend to be reported as classes rather than discrete molecules in clinical and epidemiology studies) and most endogenous molecules with large blood concentrations had few PubMed citations. Several highly-cited chemicals are familiar biomarkers of human diseases and causal exposures, e.g. cholesterol (n = 4,449, cardiovascular disease), folic acid (n = 595, cancer and neural-tube defects), lead (n = 65, cardiovascular and neurological diseases) and cotinine (n = 78, smoking-related diseases) plus vitamins, hormones and antioxidants. Aspirin was the most-cited drug (n = 515) followed by atorvastatin (n = 206).

Sequencing the human genome motivated mapping of G-centric molecular pathways at multiple levels and made metabolites with annotated pathways desirable targets for systems biology (Chen et al. 2012). When matching records from the NCBI Biosystems database were retrieved for chemicals in our inventory, at least one human metabolic pathway had been reported for 658 of them (42%). (Numbers of pathways are included in Supplemental Material, Table S1). Median numbers of pathways varied 6-fold across sources, with pollutants being significantly understudied (drugs = 4, endogenous = 6, food chemicals = 4, pollutants = 1; *P*-value < 0.0001). The chemical-similarity map of these 658 chemicals is shown in Figure 3C with symbol size representing the number of pathways. The largest numbers of pathways corresponded to purine-nucleotide phosphates (maximum = 707 for adenosine triphosphate), amino acids and derivatives, fatty acids and dietary metals. In contrast to prominent disease-risk citations that

were distributed more-or-less evenly across source categories (Figure 3B) chemicals with many pathways were overwhelmingly endogenous molecules and essential nutrients (Figure 3C).

Because the sets of PubMed and Biosystems hits were not completely overlapping, we repeated the analysis of source categories for the 267 chemicals that had at least one disease-risk citation and at least one human metabolic pathway. Results from this subset of chemicals were essentially the same as for the complete datasets. Median numbers of PubMed hits varied 2.4-fold across source categories (drugs = 7, endogenous = 7, food chemicals = 17, pollutants = 9; *P*-value = 0.0261) but did not differ significantly when food chemicals were removed (*P*-value = 0.4135). In contrast, median numbers of human-metabolic pathways varied 12-fold across source categories and were much smaller for drugs and pollutants than for endogenous and food chemicals (drugs = 4, endogenous = 11.5, food chemicals = 12, pollutants = 1; *P*-value < 0.0001).

Discussion

Discovering causes of disease

Data summarized in Figure 1 suggest that only about half of the current burden of chronic diseases can be attributed to known exposures and motivate more thorough scrutiny of the exposome to find unknown causes. This will be challenging due to the remarkable ranges of human exposures across sources and chemical classes that are displayed in Figures 2 and 3. Such extreme variation suggests that knowledge-driven studies are ill suited for discovering unknown causes of chronic diseases. There are simply too many diverse chemicals covering too great a concentration range to formulate reasonable hypotheses. We should narrow the list of chemical candidates by using EWAS to find discriminating exposures in biospecimens from diseased and healthy subjects (Holmes et al. 2008; Patel et al. 2010; Rappaport 2012; Ritchie et al. 2010; Z Wang et al. 2011), essentially following the same strategy as GWAS. Once identified, these

chemicals can be targeted to investigate sources, causality, disease mechanisms and interventions (Rappaport 2012). A good example of this two-stage strategy is given by Hazen and coworkers, who linked risks of cardiovascular disease with blood concentrations of trimethylamine-*N*-oxide, a metabolite of choline and carnitine derived from microbial/human metabolism (Koeth et al. 2013; Tang et al. 2013; Z Wang et al. 2011).

Optimally, EWAS would employ untargeted methods to compare blood exposomes between cases and controls nested in cohort studies. Although untargeted high-resolution mass spectrometry can detect more than 30,000 features of small molecules in human serum (Ivanisevic et al. 2013), use of untargeted platforms in our laboratory cannot reliably measure blood concentrations below about 0.1 µM in 50 µl of serum. Given the extraordinary dynamic range of small molecules and metals (Figure 2), this suggests that untargeted analyses will miss about 90% of pollutants and 30% of endogenous and food chemicals, including hormones (e.g. estradiol and testosterone), carcinogens (e.g., aflatoxin-B1 and benzene) and endocrine disruptors (e.g., genistein, PCBs and DDE). Thus, while increased sensitivity can be anticipated with untargeted mass spectrometry, EWAS currently require a combination of untargeted (Holmes et al. 2008; Ritchie et al. 2010; Z Wang et al. 2011) and semi-targeted (Patel et al. 2010) methods to quantify exposures. Also, as in the Human Genome Project (NHGRI 2013), different laboratories could address specific parts of the exposome in a complementary and collaborative way.

Magnitudes of exposures

Ranges of blood concentrations varied greatly within and between sources of exposure as shown in Figure 2. While we had anticipated that endogenous and food chemicals would have similar blood levels, we were surprised to observe the near-perfect overlap of concentrations of these chemicals with those of drugs. Such similar cumulative distributions suggest that blood

concentrations of endogenous human metabolites and food chemicals are in the therapeutic range of pharmacologic agents. We were also somewhat surprised to observe that blood concentrations of pollutants were 1,000 times lower than those of chemicals from other categories. Such disparate blood levels across exposure sources awaken arguments by Ames and colleagues that natural toxins and protective chemicals are consumed in much greater quantities than synthetic chemicals and, therefore, should be considered when assessing disease risks (Ames 1983; Ames et al. 1987; Ames et al. 1990a, b). This further emphasizes the importance of EWAS for interrogating all chemicals that can cause chronic diseases.

Epidemiology and systems biology

Weighting chemicals by blood concentrations (Figure 3A), epidemiologic (risk-factor) citations (Figure 3B) or human metabolic pathways (Figure 3C) altered the appearances of chemical-similarity maps. Epidemiologic citations downgraded the importance of endogenous molecules while upgrading pollutants and drugs, but weighting by numbers of metabolic pathways had the opposite effect. These markedly different maps were unanticipated because it is generally thought that epidemiology and systems biology work hand-in-glove to elucidate causes and mechanisms of disease (Nicholson et al. 2012b).

Epidemiologists are interested in causes of disease, including genetic factors (G) and exposures (E) related to metabolism, diet, pollution, infections, lifestyles and behaviors. When they have used blood concentrations to quantify chemical exposures from G, E and G×E, epidemiologists have successfully linked chronic diseases to targeted endogenous and exogenous chemicals (Figures 1 and 3B). We assumed that chemicals that had been repeatedly associated with chronic diseases (Figure 3B) would be logical candidates for exploration of metabolic pathways. However, since only 29% of the chemicals in our database with three or more PubMed risk-factor citations also had a Biosystems hit (i.e., 189/658), this was apparently not the case.

Rather, systems biologists focus on metabolic pathways that are under homeostatic control and, therefore, presume a G-centric hierarchy that culminates in the endogenous metabolome (Nicholson et al. 2012b). From the systems-biology perspective, the most meaningful metabolites are those that participate in many pathways (Loscalzo et al. 2007), and Figure 3C points to products of energy metabolism and essential nutrients as filling that role. If such molecules can be linked to disease, then their concentrations can promote early diagnosis and treatment even if causal E and G×E factors are unknown. For example, high concentrations of branched-chain amino acids (leucine, isoleucine and valine) predict incipient diabetes and offer avenues for treatment (Newgard 2012; TJ Wang et al. 2011). However, the poor track record of GWAS in explaining the variation of chronic diseases suggests that systems biologists who look only at endogenous metabolites (i.e. molecules produced under human genomic control) will miss opportunities to discover causal pathways. Indeed, only 2,626 (6.4%) (ReconX 2013) of the 41,000 small molecules currently thought to populate the human body (Wishart et al. 2013) are products of endogenous human metabolism.

The microbiome

When considering G and G×E effects it is important to remember that 90 percent of the approximately 10¹⁴ cells in the human body actually reside in the gut microbiota (Savage 1977). This superorganism contributes ~500,000 microbial protein-coding genes (Qin et al. 2010) compared to a human complement of ~20,000 protein-coding genes. Thus human biospecimens contain a plethora of bioactive molecules generated from microbial metabolism (Nicholson et al. 2012a) in addition to chemicals introduced by the diet, drugs, infectious organisms, pollution and lifestyle factors (Nicholson and Wilson 2003; Rappaport and Smith 2010). Chemicals produced by the microbiota control development and maintenance of the human immune system as well as important cell-signaling processes (Nicholson et al. 2012a) and appear to be intimately involved

in development of chronic diseases (Blumberg and Powrie 2012; Haiser and Turnbaugh 2012). Although research involving microbial contributions to the human exposome is in its infancy, it should expand dramatically as the important roles played by the microbiota are recognized in disease etiology (Koeth et al. 2013; Ridaura et al. 2013; Tang et al. 2013; Z Wang et al. 2011).

Internal and external measures of exposure

To discover unknown exposures that cause disease, we advocate data-driven EWAS to profile chemicals in blood from disease cases and controls (Rappaport 2012). Internal measures of exposure such as the blood exposome offer advantages for EWAS because they represent all sources of chemicals, including those generated inside the body, and blood specimens are often archived in prospective cohort studies (Rappaport and Smith 2010). As EWAS discover new disease associations, knowledge-driven studies will be needed to curate exposure sources and quantify exposure-response relationships - thereby strengthening causal inferences - and to suggest interventions (Rappaport 2012). To the extent that important exposures originate outside the body, this follow-up will involve exposure scientists, industrial hygienists, food scientists and analytical chemists who measure chemicals in air, water and food, as well as biologists who evaluate mechanisms of action (Lioy and Rappaport 2011; Rappaport 2011; Scalbert et al. in press; Wild 2012). Thus, the process of identifying causal exposures can require measurements of chemicals both inside and outside the body across a diverse scientific milieu.

Limitations

Because we relied on publically accessible data, our findings and their interpretation are conditioned by the chemicals compiled by HMDB and NHANES and by publications and metabolic pathways curated through NCBI. Most of the 1,561 chemicals we investigated in human blood were derived from foods and endogenous processes because these are major foci of HMDB. Most of the pollutants in our database were reported by NHANES. Yet, a roughly equal

number of other pollutants from NHANES were excluded from our database because they were not detected in most blood samples (CDC 2009, 2012, 2013). If non-detects from NHANES had been included, the shift towards lower blood concentrations of pollutants relative to chemicals from other sources would have been even greater. We also recognize that some of our data could be biased. For example, using PubMed citations to assess disease associations of particular exposures can introduce biases related to prior publications as well as research priorities for different diseases, numbers of investigators and journals, etc. As noted previously, the Biosystems database of human metabolic pathways reflects apparent biases favoring chemicals that are involved in many pathways regardless of disease associations. Finally, we were unable to investigate possible effects of chemical interactions on disease risks. But despite these limitations, the vast diversity and concentration ranges of blood chemicals should be apparent as should differences in median blood concentrations observed across source categories (Figures 2 and 3).

Conclusions

The extreme complexity and dynamic range of the blood exposome (Figures 2 and 3) should motivate data-driven studies to discover unknown causes of chronic diseases, regardless of their exogenous and endogenous origins (Rappaport 2012). Candidate exposures can be identified by EWAS that compare omic profiles in blood from diseased and healthy subjects.

The apparent disconnect between chemical-specific disease risks (Figure 3B) and human metabolic pathways (Figure 3C) indicates that systems biologists are only marginally engaged in elucidating causal disease pathways. We promote a more global approach to systems biology (Nicholson and Wilson 2003) that expands beyond the endogenous metabolome to the blood exposome, illustrated here by a large sample of circulating small molecules and inorganic species.

Perhaps the most compelling reason for embracing the blood exposome is the potential to discover all chemicals that cause disease and then to intervene so as to modify exposures and the concomitant burden of disease (Christiani 2011). The current inventory of small molecules and metals associated with chronic diseases consists of about 300 chemicals that have been targeted repeatedly in epidemiologic and clinical studies (Figure 3B). With recognition of their health significance, these chemicals have been routinely monitored for clinical interventions (e.g. cholesterol, folic acid and vitamins) and as regulated pollutants (e.g., lead, arsenic, benzene and PCBs). Yet, further scrutiny of these recognized health hazards adds little to our understanding of disease causation. If we expect to reduce the burden of chronic diseases, it is time to find the undiscovered health-impairing and health-promoting chemicals to which humans are exposed (Figure 1), not only small molecules and metals but also proteins and foreign DNA and RNA.

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Figure legends

Figure 1. Risk factors for exposures that contribute to chronic diseases. The chart was compiled from WHO estimates of exposures affecting 50 million global deaths in 2010 (Lim et al. 2012).

Figure 2. Small molecules and metals in human blood. Each curve represents the cumulative distribution of chemical concentrations from a particular source category (pollutants, n = 94; drugs, n = 49; food chemicals, n = 195; endogenous chemicals, n = 1,223). Abbreviations: OCDD, 1,2,3,4,6,7,8,9-octachlorooxanthrene; BDE 100, 2,2',4,4',6-pentabromodiphenyl ether; PCB 170, 2,2',3,3',4,4',5-heptachloro-1,1'-biphenyl; DDE, 1,1-bis-(4-chlorophenyl)-2,2-dichloroethene.

Figure 3. Chemical-similarity maps of small molecules and metals in human blood (Tanimoto coefficient ≥ 0.7 ; symbol color represents the source category).

A) All chemicals (*n* = 1,561; symbol size reflects the blood/serum concentration). Legend: AA, leucotrienes; AB, perfluorinated compounds; AC, alkylamines; AD, pteridines; AE, pyrimidine nucleotides; AF, aliphatic amino acids and derivatives; AG, sphingolipids; AH, organo-chlorine pesticides; AI, prenol lipids; AJ, sulfur compounds; AK, flavonoids; AL, pyrroles and indoles; AM, pyridines; AN, alkaloids; AO, benzoic acids and phenols; AP, eicosanoids; AQ, fatty acids and fatty amines; AR, steroids; AS, organic acids; AT, monosaccharides; AU, phosphates; AV, alcohols; AW, fatty acid esters and conjugates; AX, polychlorinated biphenyls; AY, simple aromatics; AZ, chlorinated dioxins and furans; BA, sulfates and nitrites/nitrates; BB, purine nucleotides; BC, aromatic amino acids and derivatives; BD, benzoic acids and phenols.

B) Matching chemicals from (A) cited in studies of chronic-disease risks (n = 336; symbol size reflects the number of citations). Legend: 1, Se; 2, nitric oxide; 3, folic acid; 4, vitamin B12; 5, metformin; 6, cotinine; 7, Pb; 8, bilirubin; 9, atorvastatin; 10, ascorbic acid; 11, thyroxine; 12,

norepinephrine; 13, aspirin; 14, eicosapentaenoic acid; 15, Mg; 16, Ca; 17, Na; 18, uric acid; 19, creatinine; 20, *L*-arginine; 21, homocysteine; 22, *L*-methionine; 23, *L*-valine; 24, β-carotene; 25, vitamin A; 26, vitamin D3; 27, cholesterol; 28, simvastatin; 29, aldosterone; 30, cortisol; 31, testosterone; 32, malondialdehyde; 33, *D*-glucose; 34, estradiol; 35, PCBs; 36, ethanol.

C) Matching chemicals from (A) having human metabolic pathways (*n* = 658; symbol size reflects the number of pathways). Legend: 1, adenosine triphosphate; 2, hydrogen peroxide; 3, adenosine diphosphate; 4, guanosine diphosphate; 5, guanosine triphosphate; 6, NADPH; 7, cyclic AMP; 8, adenosine monophosphate; 9, NADH; 10, NAD; 11, FAD; 12, Mn; 13, Na; 14, Ca; 15, Zn; 16, Mg; 17, K; 18, norepinephrine; 19, epinephrine; 20, *L*-phenylalanine; 21, *L*-tyrosine; 22, dopamine; 23, palmitic acid; 24, cholesterol; 25, *L*-glutamic acid; 26, adenine; 27, *L*-aspartic acid; 28, oxoglutaric acid; 29, pyruvic acid; 30, phosphate; 31, pyrophosphate; 32, formic acid; 33, uridine 5'-monophosphate; 34, uridine 5'-diphosphate; 35, *L*-arginine; 36, *L*-alanine; 37, *L*-cysteine; 38, *L*-serine; 39,arachodonic acid; 40, α-linolenic acid.

Figure 1.

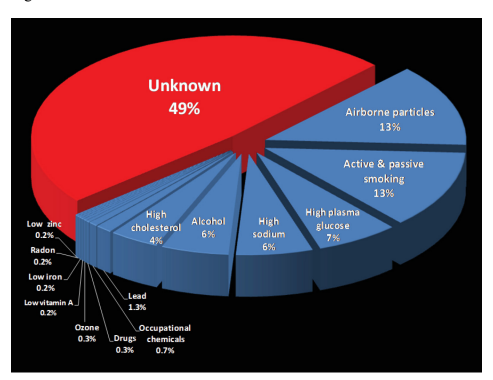


Figure 2.

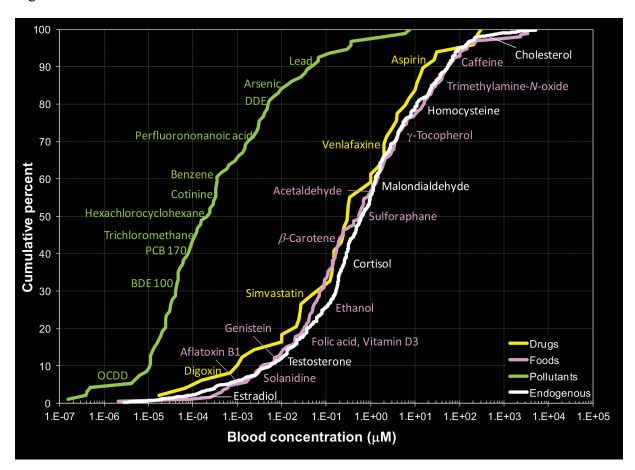
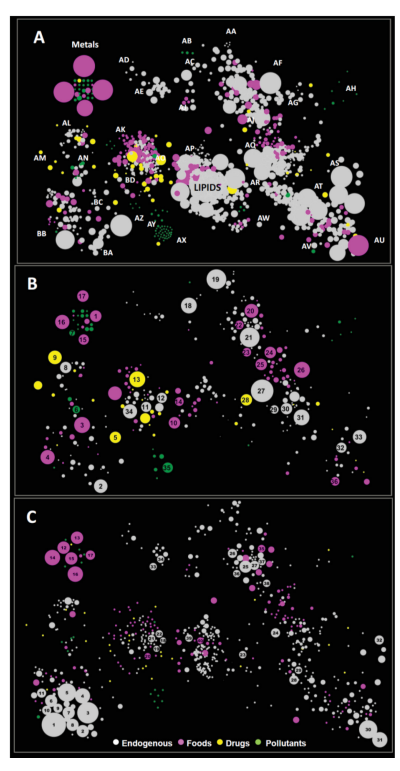


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